L Number	Hits	Search Text	DB	Time stamp
9	119	flavivirus and vero	USPAT;	2002/02/19 15:05
			US-PGPUB;	
			EPO; JPO;	
			DERWENT;	
			IBM_TDB	
10	13555	(flavivirus and vero) and yellow fever	USPAT;	2002/02/19 15:05
			US-PGPUB;	
			EPO; JPO;	·
			DERWENT;	
			IBM_TDB	
11	119	(,	USPAT;	2002/02/19 15:06
		virus)	US-PGPUB;	<i>j</i>
			EPO; JPO;	! !
			DERWENT;	
			IBM_TDB	0000/00/10 15 10
12	92	1,	USPAT;	2002/02/19 15:19
		virus)) and vaccine	US-PGPUB;	
			EPO; JPO;	
			DERWENT;	
1 2 2		(//51	IBM_TDB	2002/02/19 15:19
13	2	(((flavivirus and vero) and (yellow fever	USPAT;	2002/02/19 15:19
		virus)) and vaccine) and yf17d	US-PGPUB;	
			EPO; JPO;	
			DERWENT; IBM TDB	
			I TOM_TOB	

Search History

ΤI Preparation of inactivated viral vaccines Wiesehahn, Gary P., Alameda, CA, United States TN Creagan, Richard P., Alta Loma, CA, United States Stevens, David R., Fremont, CA, United States Giles, Richard, Alameda, CA, United States AB Vaccines employing inactivated viruses having improved retention of antigenic characteristics are prepared by psoralen-inactivation of the live virus in a non-oxidizing atmosphere. By excluding oxygen and other oxidizing species from the inactivation medium, degradation of the antigen characteristics resulting from irradiation with ultraviolet light is largely prevented. The resulting inactivated viruses are employed in vaccine preparations for the inoculation of susceptible hosts to inhibit viral infection. 87:65285 USPATFULL ΑN ΤI Preparation of inactivated viral vaccines IN Wiesehahn, Gary P., Alameda, CA, United States Creagan, Richard P., Alta Loma, CA, United States Stevens, David R., Fremont, CA, United States Giles, Richard, Alameda, CA, United States PA Advanced Genetics Research Institute, Oakland, CA, United States (U.S. corporation) US 4693981 19870915 <--PΙ ΑI US 1985-785354 19851007 (6) DCD 20021008 Continuation-in-part of Ser. No. US 1983-563939, filed on 20 Dec 1983, RLI now patented, Pat. No. US 4545987 And a continuation-in-part of Ser. No. US 1984-592661, filed on 23 Mar 1984, now abandoned DTUtility Primary Examiner: Rose, Shep K. EXNAM Rowland, Bertram I. LREP Number of Claims: 9 CLMN Exemplary Claim: 1 ECL DRWN No Drawings LN.CNT 1219

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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ANSWER 38 OF 81 USPATFULL
L7
       Flavivirus recombinant poxvirus vaccine
TI
       Paoletti, Enzo, Delmar, NY, United States
IN
       Pincus, Steven E., East Greenbush, NY, United States
       What is described is a recombinant poxvirus, such as vaccinia virus,
AΒ
       fowlpox virus and canarypox virus, containing foreign DNA from
       flavivirus, such as Japanese encephalitis virus, yellow
     fever virus and Dengue virus. In a preferred
       embodiment, the recombinant poxvirus generates an extracellular
particle
       containing flavivirus E and M proteins capable of inducing neutralizing
       antibodies, hemagglutination-inhibiting antibodies and protective
       immunity against flavivirus infection. What is also described is a
       vaccine containing the recombinant poxvirus for inducing an
       immunological response in a host animal inoculated with the vaccine.
       96:38606 USPATFULL
AN
ΤI
       Flavivirus recombinant poxvirus vaccine
       Paoletti, Enzo, Delmar, NY, United States
IN
       Pincus, Steven E., East Greenbush, NY, United States
PΑ
       Virogenetics Corporation, Troy, NY, United States (U.S. corporation)
PΙ
       US 5514375 19960507
ΑI
       US 1991-714687 19910613 (7)
       Continuation-in-part of Ser. No. US 1991-711429, filed on 6 Jun 1991,
RLI
       now abandoned And a continuation-in-part of Ser. No. US 1991-713967,
       filed on 11 Jun 1991, now abandoned which is a continuation-in-part of
       Ser. No. US 1991-666056, filed on 7 Mar 1991, now abandoned , said Ser.
                -711429 which is a continuation of Ser. No. US 1990-567960,
       filed on 15 Aug 1990, now abandoned
DT
       Utility
EXNAM Primary Examiner: Sidberry, Hazel F.; Assistant Examiner: Tuscan,
       Michael
       Curtis, Morris & Safford
LREP
       Number of Claims: 22
CLMN
       Exemplary Claim: 1
ECL
       19 Drawing Figure(s); 16 Drawing Page(s)
DRWN
LN.CNT 2530
CAS INDEXIN
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- L6 ANSWER 8 OF 17 MEDLINE
- TI Attenuation of wild-type yellow fever virus by passage in HeLa cells.
- AU Barrett A D; Monath T P; Cropp C B; Adkins J A; Ledger T N; Gould E A; Schlesinger J J; Kinney R M; Trent D W
- AB During the 1960s three different research groups reported that passage of wild-type yellow fever (YF) virus [strain Asibi (YF-Asibi)] in HeLa cells resulted in attenuation of the virus for monkeys so that the virus no longer caused viscerotropic disease. We have repeated and extended this observation to analyse the process of attenuation of YF virus during cell culture passage. A large plaque (LP) variant of YF-Asibi virus became attenuated for both monkeys and mice following six serial subcultures in HeLa cells (YF-Asibi-LP HeLa p6). Thus, attenuation was probably due to a genetic change in the virus population rather than to selective

## enrichment

of a pre-existing variant of YF-Asibi-LP virus. No evidence was obtained to implicate defective interfering particles in the attenuation process. Comparison of the YF-Asibi-LP viruses before and after passage in HeLa cells, using a panel of envelope protein-reactive monoclonal antibodies (MAbs), showed that MAbs which specifically neutralize YF-Asibi-LP virus, and not YF 17D-204 vaccine virus, also neutralized

YF-Asibi-LP HeLa p6. This indicated that the epitopes involved in the biological process of neutralization were not altered during attenuation. However, two MAbs that recognize envelope protein epitopes did

distinguish

YF-Asibi-LP

between HeLa- and non-HeLa-passaged YF-Asibi-LP virus. One of these (MAb 117) which is YF wild-type-specific, recognized YF-Asibi-LP virus but not YF-Asibi-LP HeLa p6 virus, whereas the other (MAb411), which is YF vaccine-specific, recognized YF-Asibi-LP HeLa p6 virus but not

virus. These results suggest that antigenic changes in the viral envelope protein may determine the relative virulence or attenuation of YF virus.

- AN 91037961 MEDLINE
- DN 91037961 PubMed ID: 2230735
- TI Attenuation of wild-type yellow fever virus by passage in HeLa cells.
- AU Barrett A D; Monath T P; Cropp C B; Adkins J A; Ledger T N; Gould E A; Schlesinger J J; Kinney R M; Trent D W
- CS Department of Microbiology, University of Surrey, Guildford, U.K.
- SO JOURNAL OF GENERAL VIROLOGY, (1990 Oct) 71 ( Pt 10) 2301-6. Journal code: I9B; 0077340. ISSN: 0022-1317.
- CY ENGLAND: United Kingdom
- DT Journal; Article; (JOURNAL ARTICLE)
- LA English
- FS Priority Journals
- EM 199012
- ED Entered STN: 19910208

Last Updated on STN: 19970203 Entered Medline: 19901204

QR1. J6

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L6 ANSWER 11 OF 17 MEDLINE
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TI Conditions for haemolysis by flaviviruses and characterization of the haemolysin.

AU Cammack N; Gould E A

AB The 17D vaccine strain of yellow fever virus (YF 17D) was used to establish the optimal conditions for lysis of chick erythrocytes. Tissue culture-grown, polyethylene glycol-concentrated virus

showed peak activity at pH 5.4 in citrate buffer when incubated at 37 degrees C. A further two- to fourfold increase in titre was obtained by pretreatment of the chick erythrocytes with 250 micrograms/ml trypsin. These conditions were also shown to be optimal for Japanese encephalitis (JE), West Nile (WN) and dengue-2 (den2) viruses. The ratio of haemagglutination (HA) titre to haemolysis (HL) titre approximated to unity, suggesting that the two functions are associated with the same molecule although as separable entities since selective inactivation of the HL activity of the virus was accomplished using 60 micrograms/ml trypsin. HL could be demonstrated at neutral pH if the chick erythrocytes were first subjected to treatment with acidic pH buffer. The effect on

the

virus envelope is thus not the sole contribution of a low pH environment to optimal HL. Hyperimmune rabbit antiserum prepared against purified YF 17D virions inhibited HA and HL if added before

agglutination had occurred by the virus but when added after agglutination

had taken place it showed specific anti-HL activity. Monoclonal antibodies

that inhibited HA (HAI) by YF 17D did not inhibit HL

(HLI) activity when applied after agglutination had taken place.

## Moreover,

monoclonal antibodies specific for the 54K glycoprotein of YF virus but without HAI activity also had no effect on HL when added either before or after agglutination. As yet, we have been unable to identify a monoclonal antibody displaying specific anti-HL activity but all those directed against the 54K envelope glycoprotein possessing HAI activity showed HA

to

be a prerequisite for HL.

AN 86010270 MEDLINE

DN 86010270 PubMed ID: 2995565

TI Conditions for haemolysis by flaviviruses and characterization of the haemolysin.

AU Cammack N; Gould E A

SO JOURNAL OF GENERAL VIROLOGY, (1985 oct) 66 ( Pt 10) 2291-6. Journal code: I9B; 0077340. ISSN: 0022-1317.

CY ENGLAND: United Kingdom

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 198511

Last Updated on STN: 19900321 Entered Medline: 19851108

- ANSWER 9 OF 81 MEDLINE L7
- Virulence and pathogenesis of yellow fever ΤI
- virus serially passaged in cell culture.
  Converse J L; Kovatch R M; Pulliam J D; Nagle S C Jr; Snyder E M ΑU
- ΑN 71267151 MEDLINE
- PubMed ID: 4998347 DN 71267151
- Virulence and pathogenesis of yellow fever virus serially passaged in cell culture. ΤI
- Converse J L; Kovatch R M; Pulliam J D; Nagle S C Jr; Snyder E M ΑU
- APPLIED MICROBIOLOGY, (1971 Jun) 21 (6) 1053-7. SO Journal code: 6K0; 7605802. ISSN: 0003-6919.
- CY United States
- Journal; Article; (JOURNAL ARTICLE) DT
- LΑ English
- FS Priority Journals
- EM 197110
- Entered STN: 19900101 ED

Last Updated on STN: 19970203 Entered Medline: 19711015

- L7 ANSWER 6 OF 7 MEDLINE
- TI THE GROWTH OF ASIBI STRAIN YELLOW FEVER VIRUS IN TISSUE CULTURES. II. MODIFICATION OF VIRUS AND CELLS.
- AU HARDY F M
- AN 64001668 MEDLINE
- DN 64001668
- TI THE GROWTH OF ASIBI STRAIN YELLOW FEVER
  VIRUS IN TISSUE CULTURES. II. MODIFICATION OF VIRUS AND CELLS.
- AU HARDY F M
- SO JOURNAL OF INFECTIOUS DISEASES, (1963 JUL-AUG) 113 9-14. Journal code: IH3. ISSN: 0022-1899.
- CY United States
- DT Journal
- LA English
- FS OLDMEDLINE
- EM 196401
- ED Entered STN: 19990716

Last U

- L7 ANSWER 5 OF 7 MEDLINE
- TI GROWTH OF 17D YELLOW FEVER VIRUS AND FACTORS INFLUENCING ITS TRANSMISSION WITHIN CELL CULTURES IN VITRO.
- AU LITWIN J
- AN 64143518 MEDLINE
- DN 64143518
- TI GROWTH OF 17D YELLOW FEVER VIRUS AND FACTORS INFLUENCING ITS TRANSMISSION WITHIN CELL CULTURES IN VITRO.
- AU LITWIN 3
- SO ACTA PATHOLOGICA ET MICROBIOLOGICA SCANDINAVICA, (1964) 61 605-18. Journal code: 100. ISSN: 0365-5555.
- CY Denmark
- DT Journal
- LA English
- FS OLDMEDLINE
- EM 196412
- ED Entere

ANSWER 2 OF 7 MEDLINE DUPLICATE 2 L7 Growth of 17D yellow fever virus in a TImacrophage-like cell line, U937: role of Fc and viral receptors in antibody-mediated infection. ΑIJ Schlesinger J J; Brandriss M W Growth characteristics of 17D yellow fever virus (17D-YF) and conditions for infection were studied in U937, a macrophage-like, Fc receptor-bearing continuous human cell line. Antibody to 17D-YF was obtained by immunization of normal subjects with 17D-YF vaccine. Cells were infected in the presence or absence of immune whole sera or immunoglobulin fractions. Infection of U937 was temperature dependent; the yield of virus was variable but at low temperature viral titers were consistently higher when infection was established in the presence of antibody. Results of infectious center assays indicated that the increased yield of virus was largely or entirely due to an increase in the number of cells producing virus early in the course of infection. Enhancement of viral growth was mediated by IgG but not IgM fractions of immune sera. Trypsinization of U937 resulted in a 90 to 95% reduction of infection in the absence of antibody but in the presence of antibody viral titers were higher in trypsinized than in nontrypsinized cells. Antibody to 17D-YF, contained in the whole IgG fraction of sera, bound to U937 to mediate infection without first being complexed to virus. Preincubation of U937 with IgG1 but not IgG2 myeloma proteins abrogated antibody-mediated infection. This result is compatible with the known affinities of U937 Fc receptors for specific subclasses of IgG and provides evidence for the role of the Fc receptor in antibody-mediated enhancement of viral growth. Persistent infection characterized by a lack of detectable cytopathogenic effect was established in long-term cultures of U937. This pattern of infection might be related to the unique effectiveness of the 17D-YF vaccine. MEDLINE 81240805 AN PubMed ID: 7252155 DN 81240805 TIV. Growth of 17D yellow fever virus in a macrophage-like cell line, U937: role of Fc and viral receptors in antibody-mediated infection. ΑU Schlesinger J J; Brandriss M W JOURNAL OF IMMUNOLOGY, (1981 Aug) 127 (2) 659-65. SO Journal code: IFB; 2985117R. ISSN: 0022-1767. United States CYQR 180. JG Journal; Article; (JOURNAL ARTICLE) DT

Abridged Index Medicus Journals; Priority Journals

LΑ

FS EM

ED

English

198109

Entered STN: 19900316

Last Updated on STN: 19900316 Entered Medline: 19810915